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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/657,404	09/08/2003	Gavin William Halbert	031749/268956	3177
826 7590 10/02/2007 ALSTON & BIRD LLP BANK OF AMERICA PLAZA			EXAMINER	
			ROBINSON, HOPE A	
101 SOUTH TRYON STREET, SUITE 4000 CHARLOTTE, NC 28280-4000		4000	ART UNIT	PAPER NUMBER
	,		1652	
			WALL DATE	DEL IVERY MODE
			MAIL DATE	DELIVERY MODE
			10/02/2007	PAPER

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

	•	Application No.	Applicant(s)				
•		10/657,404	HALBERT ET AL.				
	Office Action Summary	Examiner	Art Unit				
	·	Hope A. Robinson	1652				
Period fo	The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply						
WHIC - Exte after - If NC - Failu Any	ORTENED STATUTORY PERIOD FOR REPLY CHEVER IS LONGER, FROM THE MAILING DANSIONS of time may be available under the provisions of 37 CFR 1.13 SIX (6) MONTHS from the mailing date of this communication. Operiod for reply is specified above, the maximum statutory period we are to reply within the set or extended period for reply will, by statute, reply received by the Office later than three months after the mailing ed patent term adjustment. See 37 CFR 1.704(b).	ATE OF THIS COMMUNICATION 16(a). In no event, however, may a reply be still apply and will expire SIX (6) MONTHS from cause the application to become ABANDO	ON. timely filed om the mailing date of this communication. NED (35 U.S.C. § 133).				
Status							
1)⊠	Responsive to communication(s) filed on 13 Ja	<u>nuary 2007</u> .					
,	This action is FINAL . 2b) This action is non-final.						
3)	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is						
closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.							
Disposit	ion of Claims						
4)⊠	4)⊠ Claim(s) <u>1,2,4-10 and 12-25</u> is/are pending in the application.						
	4a) Of the above claim(s) <u>16-25</u> is/are withdrawn from consideration.						
·	5) Claim(s) is/are allowed.						
· <u> </u>	s)⊠ Claim(s) <u>1,2,4-10,12 and 13</u> is/are rejected.						
•	Claim(s) <u>14 and 15</u> is/are objected to.						
8)[_]	8) Claim(s) are subject to restriction and/or election requirement.						
Applicati	ion Papers						
9)	The specification is objected to by the Examiner						
10) ☐ The drawing(s) filed on is/are: a) ☐ accepted or b) ☐ objected to by the Examiner.							
Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a).							
Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).							
11) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152.							
Priority u	under 35 U.S.C. § 119						
	Acknowledgment is made of a claim for foreign All b) Some * c) None of:	priority under 35 U.S.C. § 119(a)-(d) or (f).				
1. Certified copies of the priority documents have been received.							
2. Certified copies of the priority documents have been received in Application No							
3. Copies of the certified copies of the priority documents have been received in this National Stage							
application from the International Bureau (PCT Rule 17.2(a)).							
* See the attached detailed Office action for a list of the certified copies not received.							
Attachmen	t(s) ·						
_	e of References Cited (PTO-892)	4) Interview Summa					
2) Notic	e of Draftsperson's Patent Drawing Review (PTO-948)	Paper No(s)/Mail 5) Notice of Informal	Date				
	mation Disclosure Statement(s) (PTO/SB/08) r No(s)/Mail Date	6) Other:	т акти друговноги				

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DETAILED ACTION

Application Status

1. A request for continued examination under 37 CFR 1.114, including the fee set forth in 37 CFR 1.17(e), was filed in this application after final rejection. Since this application is eligible for continued examination under 37 CFR 1.114, and the fee set forth in 37 CFR 1.17(e) has been timely paid, the finality of the previous Office action has been withdrawn pursuant to 37 CFR 1.114. Applicant's submission filed on July 13, 2007, has been entered.

Claim Disposition

2. Claims 1-2, 4-10 and 12-25 are pending. Claims 1-2 and 4-10 and 12-15 are under examination.

New-Claim Objection

3. Claim 15 is objected to because of the following informalities:

For clarity, precision and consistency of claim language it is suggested that claim15 is amended to recite, "The particle according to claim 1", see for example claim 8.

Correction is required.

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The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

4. Claims 1-2 and 4-10 and 12-13, are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention.

The claimed invention is directed to a non-naturally occurring receptor low density lipoprotein particle comprising at least one peptide component, said peptide component comprises at least a binding site for an Apo B protein receptor and wherein the binding sequence of the peptide component has at least 70% amino acid sequence identity to the amino acid selected from the group consisting of SEQ ID NO:1, 2 and 8. In addition, some claims are directed to 80% or 90% sequence identity. Thus the claims are directed to a genus that is highly variable. For example, the recited 70% sequence identity to SEQ ID NO:2 which comprises 11 residues would mean approximately 8 residues through the sequence could be deleted, inserted, substituted, etc. The specification fails to provide any additional representative species of the claimed genus to show that applicant was in possession of the claimed genus. A

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representative number of species means that the species, which are adequately described are representative of the entire genus. The written description requirement for a claimed genus may be satisfied through sufficient description of a representative number of species by actual reduction to practice, disclosure of drawings, or by disclosure of relevant identifying characteristics, for example, structure or other physical and/or chemical properties, by functional characteristics coupled with a known or disclosed correlation between function and structure, or by a combination of such identifying characteristics, sufficient to show the applicant was in possession of the claimed genus.

Accordingly, in the absence of sufficient recitation of distinguishing identifying characteristics, the specification does not provide adequate written description of the claimed genus. Therefore, the genus of claimed polypeptides encompasses widely variant species. Further, *Vas-Cath Inc. v. Mahurkar*, 935 F.2d 1555, 1563-64, 19 USPQ2d 1111, 1117 (Fed. Cir.1991), states that "applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in *possession of the invention*. The invention is, for purposes of the 'written description' inquiry, *whatever is now claimed*" (See page 1117). The specification does not "clearly allow persons of ordinary skill in the art to recognize that [he or she] invented what is claimed" (See *Vas-Cath* at page 1116). The skilled artisan cannot envision the detailed chemical structure of the encompassed genus of polypeptides, and therefore, conception is not achieved until reduction to practice has occurred, regardless of the complexity or simplicity of the method of isolation. Adequate written

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description requires more than a mere statement that it is part of the invention and reference to a potential method of isolating it. The compound itself is required. See Fiers v. Revel, 25 USPQ2d 1601 at 1606 (CAFC 1993).

Therefore, for all these reasons the specification lacks adequate written description, and one of skill in the art cannot reasonably conclude that the applicant had possession of the claimed invention at the time the instant application was filed.

5. Claims 1-2 and 4-13 are rejected under 35 U.S.C. 112, first paragraph, because the specification, while being enabling for the peptides set forth in SEQ ID NOS: 3, 4, 5, 6, 7 and 9 does not reasonably provide enablement for any peptide fragment thereof. The specification does not enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make the invention commensurate in scope with these claims. The enablement requirement refers to the requirement that the specification describe how to make and how to use the invention. There are many factors to be considered when determining whether there is sufficient evidence to support a determination that a disclosure does not satisfy the enablement requirement and whether any necessary experimentation is undue. These factors include, but are not limited to: Quantity of Experimentation Necessary; Amount of direction or guidance presented; Presence or absence of working examples; Nature of the Invention; State of the prior art and Relative skill of those in the art; Predictability or unpredictability of the art and Breadth of the claims (see In re Wands, 858 F.2d at 737, 8 USPQ2d at1404 (Fed. Cir. 1988). The factors most relevant to the instant invention are discussed below.

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The amount of experimentation required to practice the claimed invention is undue as the claims encompass an unspecified amount of peptide components (see claim 1 for example). No correlation is made between function and structure. The claims broadly read on any peptide component comprising at least one binding site for an Apo B protein receptor wherein the binding sequence of the peptide component is at least 70%, 80% or 90% identical to any Apo B protein binding sequence. Therefore, the claims encompass a genus of peptides which are needed demonstrated or described in the instant specification. A skilled artisan would recognize the unpredictability of testing the large variable genus of peptides encompassed in the claims with the expectation of finding ones that have the desired properties. This is merely an invitation to the skilled artisan to use the current invention as a starting point for further experimentation. Thus, undue experimentation would be required for a skilled artisan to make and/or use the claimed invention commensurate in scope with the claims.

Predictability of which potential changes can be tolerated in a protein's amino acid sequence and obtain the desired activity requires a knowledge of and guidance with regard to which amino acids in the protein's sequence, if any, are tolerant of modification and which are conserved (for example, expectedly intolerant to modification), and detailed knowledge of the ways in which the protein's structure relates to its function. The state of the prior art provides evidence for the high degree of unpredictability as stated above. While recombinant and mutagenesis techniques are known in the art, it is not routine in the art to screen large numbers of mutated proteins where the expectation of obtaining similar activity is unpredictable based on the instant

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disclosure. The amino acid sequence of a protein determines its structural and functional properties, and predictability of what mutations can be tolerated in a protein's sequence and result in certain activity, which is very complex, and well outside the realm of routine experimentation, because accurate predictions of a protein's function from mere sequence data are limited, therefore, the general knowledge and skill in the art is not sufficient, thus the specification needs to provide an enabling disclosure.

The working examples provided do not rectify the missing information in the instant specification pertaining to the claimed variant. The nature and properties of this claim is difficult to ascertain from the examples provided as one of skill in the art would have to engage in undue experimentation to construct the variants of the claimed invention and examine the same for function. Thus the specification does not provide support for the broad scope of the claims. The issue in this case is the breath of the claims in light of the predictability of the art as determined by the number of working examples, the skill level artisan and the guidance presented in the instant specification and the prior art of record. This make and test position is inconsistent with the decisions of In re Fisher, 427 F.2d 833, 839, 166 USPQ 18, 24 (CCPA 1970) where it is stated that "...scope of claims must bear a reasonable correlation to scope of enablement provided by the specification to persons of ordinary skill in the art...". Without sufficient guidance, determination of having the desired biological characteristics is unpredictable and the experimentation left to those skilled in the art is unnecessarily and improperly extensive and undue. See In re Wands, 858 F.2d at 737, 8 USPQ2d at 1404 (Fed. Cir. 1988). Thus, for all these reasons, the specification is not

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considered to be enabling for one skilled in the art to make and use the claimed invention as the amount of experimentation required is undue.

Response to Arguments

6. The response filed have has been considered. Note that the rejections under 35 U.S.C. 112, first paragraphs for the reasons stated above and herein, remains. Applicant state that the claims have been amended to remove recitation of "dimers thereof or analogs thereof" and to incorporate the percent language in the claim. However the claims remain problematic because the amendatory language imports an unspecified amount of variants/fragments into the claims. The claims do not identify any conserved regions or identify where in the sequence modifications will occur. Thus no correlation is made between structure and function for all the fragments/variants encompassed in the claims. Thus, the rejections of written description (lacking description) and enablement (lacking enablement) remain.

Conclusion

7. No claims are allowable as claim 14-15 are objected to as depending from a rejected base claim.

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8. Applicant's amendment necessitated the new/modified ground(s) of rejection presented in this Office action. Accordingly, **THIS ACTION IS MADE FINAL**. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the date of this final action.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Hope A. Robinson whose telephone number is 571-272-0957. The examiner can normally be reached on Monday-Friday.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Ponnathapu Achutamurthy, can be reached at (571) 272-0928. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

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Business Center (EBC) at 866-217-9197 (toll-free).

Hope Robinson, MS

Primary Examiner

HOPE ROBINSON PRIMARY EXAMINER